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⑱ Benannte Vertragsstaaten:
AT BE CH DE DK ES FR GB GR IE IT LI LU NL
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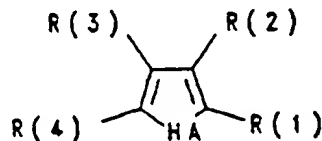
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㉑ Substituierte N-Heteroaroylguanidine, als Inhibitoren des zellulären Natrium-Protonen-Antip
als Antiarrhythmika und als Inhibitoren der Proliferation von Zellen.

㉒ Die Erfindung betrifft Heteroaroylguanidine der Formel I



worin die Substituenten HA und R(1) bis R(5) die in Anspruch 1 wiedergegebenen Bedeutungen haben. Diese Verbindungen I haben sehr gute antiarrhythmische Eigenschaften aufweisen, wie sie zum Behar Krankheiten wichtig sind, die beispielsweise bei Sauerstoffmangelerscheinungen auftreten. Die Verb sind infolge ihrer pharmakologischen Eigenschaften als antiarrhythmische Arzneimittel mit cardiop Komponente zur Infarktprophylaxe und der Infarktbehandlung sowie zur Behandlung der angina hervorragend geeignet, wobei sie auch präventiv die pathophysiologischen Vorgänge beim Entsteh misch induzierter Schäden, insbesondere bei der Auslösung ischämisch induzierter Herzarrhythmien, i oder stark vermindern. Wegen ihrer schützenden Wirkungen gegen pathologische hypoxische und iscl Situationen können die erfindungsgemäßen Verbindungen der Formel I infolge Inhibition des zelluläre Austauschmechanismus als Arzneimittel zur Behandlung aller akuten oder chronischen durch Ischämie

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Number of Countries: 022 Number of Patents: 009

Patent Family:

| Patent No | Kind | Date | Applicat No | Kind | Date | Main IPC | Week |
|------------|------|----------|-------------|------|----------|--------------|----------|
| EP 676395 | A2 | 19951011 | EP 95105088 | A | 19950405 | C07D-207/40 | 199546 B |
| DE 4412334 | A1 | 19951019 | DE 4412334 | A | 19940411 | C07D-207/416 | 199547 |
| AU 9516354 | A | 19951019 | AU 9516354 | A | 19950407 | C07D-207/416 | 199549 |
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| EP 676395 | A3 | 19960306 | EP 95105088 | A | 19950405 | C07D-207/40 | 199624 |

Priority Applications (No Kind Date): DE 4412334 A 19940411

Cited Patents: No search report pub.; 3. journal ref.; DE 1965267; DE 2055727; EP 416499; EP 556672; EP 556673; EP 556674; EP 577024; EP 589336; EP 590455; EP 622356; EP 639573; JP 44030268; WO 9304048

Patent Details:

| Patent | Kind | Lan | Pg | Filing Notes | Application | Patent |
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| DE 4412334 | A1 | 41 |
| JP 7291927 | A | 28 |
| ZA 9502930 | A | 70 |

Abstract (Basic): EP 676395 A

Heteroaryl-guanidine derivs. of formula (I) and their salts are new. A = S(O)m, O or NR5; m = 0, 1 or 2; R5 = H, 1-8C alkyl or CmH2mR81; R81 = 3-8C cycloalkyl, phenyl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe, and NR82R83) or 1-9C heteroaryl (bonded via C or N and opt. substd. by 1-3 of F, Cl, CF3, Me, OMe, OH, NH2, NHMe and NMe2); R82, R83 = H or Me; one of R1, R2 = CO-N=C(NH2)2; the other = H, F, Cl, Br, I, 1-3C alkyl, OR6, 1-4C perfluoroalkyl, CO-N=C(NH2)2 or NR6R7; R6, R7 = H or 1-3C alkyl; R3, R4 = (i) H, F, Cl, Br, I, CN, X(CH2)m(1-6C) perfluoroalkyl, X(CH2)mF, S(O)mR8, CONR9R10, COR11, SO2NR12R13; (ii) 1-8C alkyl, CmH2mR81; (iii) 1-9C heteroaryl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe, OH, NH2, NHMe and NMe2); (iv) -Y-C6H4-(CO)i-(CHOH)j-(CHOH)k-R23; (v) H, F, Cl, Br, I, CN, 1-8C alkyl, 1-8C perfluoroalkyl, 3-8C alkenyl, CgH2g-R26; SR29, OR30, NR31R32, CR33R34R35; (vii) -W-C6H4-R97; (viii) S(O)mR37, SO2NR38R39; (ix) X1R46; (x) SR64, OR65, NHR66, NR67R68, CHRR69R70, CR54R55-OH, Ctriple bondC-R56 C(R58) C-R57 (sic), (CR59R60)u-CO-(CR61R62)v-R63; (xi) SO2NHR76; or (xii) NR84R85; X = O, S or NR14; R14 = H or 1-3C alkyl; R8 = 1-5C alkyl, 3-6C alkenyl, CnH2nR15 or CF3; R9, R11, R12 = H or as R8; n = 0-4; R15 = 3-7C cycloalkyl or phenyl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe and NR16R17); R16, R17 = H or 1-4C alkyl; R10, R13 = H or 1-4C alkyl; or R9+R10 or R12+R13 = (CH2)4 or (CH2)5 in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl; R18 = 3-8C cycloalkyl or phenyl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe and NR19R20); R19, R20 = H or Me; Y = O, S or NR22; h = 0 or 1; i, j, k = 0-4; provided that h, i and k are not all 0; R22, R23 = H or 1-3C alkyl; g = 0-4; R26 = 3-8C cycloalkyl, phenyl, biphenyl, or naphthyl (where aromatics are opt. substd. by 1-3 of F, Cl, CF3, Me, OMe and NR27R28); R27, R28 = H, 1-4C alkyl or 1-4C perfluoroalkyl; R29-R31, R33 = -(CH2)m- (1-9C) heteroaryl (opt. substd. as in R81); R32, R34, R35 = H, 1-4C alkyl, 1-4C perfluoroalkyl, or as R29; R96 = heteroaryl as defined for R81, or benzyl; W = O, S or NR36; R36 = H or 1-4C alkyl; R37 = 1-8C alkyl, 1-8C perfluoroalkyl, 3-8C alkenyl or -Csh2s-R40; s = 0-4; R40 = as R26; R38 = H, 1-8C alkyl, 1-8C perfluoroalkyl, 3-8C alkenyl or -Cwh2w-R26; R39 = H, 1-4C alkyl or 1-4C perfluoroalkyl; or R38+R39 = (CH2)4 or (CH2)5, in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl; X1 = O, S, NR47, (D=O)A'- or NR48C=MN*(R49)-; M = O or S; A' = O or NR50; D = C or SO; R46, R49 = 1-8C alkyl, 3-8C alkenyl, -(CH2)b-(1-7C)perfluoroalkyl or -CxH2x-R26; b = 0 or 1; x = 0-4; R47, R48, R50 = H, 1-4C alkyl or 1-4C perfluoroalkyl; or R46+R47 or R46+R48 = (CH2)4 or (CH2)5 in which CH2 may be replaced by O, S, NH, NMe or N-benzyl; A' and N* are bonded to the phenyl ring of the benzoylguanidine structure; R64-R67, R69 = -(CH2)y-(CHOH)z-(CH2)q'-(CH2OH)t-R71 or -(CH2)b'-O-(CH2CH2O)c'-R72; R71,

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R72 = H or Me; b', c' are not defined; u, t = 1-4; v, y, z, a' = 0-4; R68, R70, R54, R55 = H or 1-6C alkyl; or CR69R70 or CR54R55 = 3-8C cycloalkylidene; R63 = H, 1-6C alkyl, 3-8C cycloalkyl or -CeH2e-R73; e = 0-4; R80 = 5-7C cycloalkyl or phenyl (opt. substd. by 1-3 of F, Cl, CF3, OMe and 1-4C alkyl); or R77+R78 = (CH2)4 or (CH2)5, in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl; R79 = as R77; or amidino; R84, R85 = H or 1-4C alkyl; or R84+R85 = (CH2)4 or (CH2)5 in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl or 1 or 2 CH2 gps. may be replaced by CH-Cd'H2d'+1; d' is not defined. Cpds. (I; A = O; R1 = -CON=C(NH2)2; R2, R3 = H; R4 = H, Me or Et) are excluded.

USE - (I) are used for treatment of arrhythmia or shock states; for treatment or prophylaxis of cardiac infarct, angina pectoris, cardiac ischaemia, ischaemic states of the peripheral and central nervous system, stroke or ischaemic states of the peripheral organs and limbs; and adjuvant during surgical operations and organ transplants; in preservation and storage of transplants; for treatment of diseases in which cell proliferation is a prim. or sec. cause, esp. atherosclerosis, complications following diabetes, cancer, fibrotic diseases, (e.g. fibrosis of the lungs, liver or kidneys) or prostatic hyperplasia; and as reagents for inhibiting Na+/H+ exchange and for diagnosis of hypertension and proliferative diseases (all claimed). More generally (I) inhibit the cellular Na+/H+ exchange mechanism and cell proliferation and are useful for combatting oxygen deficiency states, pathological hypoxia and ischaemia. They are esp. useful as antiarrhythmic agents.

Daily dose is 0.001-10 (pref. 0.01-1) mg orally, parenterally, rectally or by inhalation.

ADVANTAGE - (I) have good antiarrhythmic activity, without undesirable salidiuretic side effects, potent cellular Na+/H+ exchange inhibiting activity and good water solubility (facilitating i.v. admin.).

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Abstract (Equivalent): DE 4412334 A

Heteroaryl-guanidine derivs. of formula (I) and their salts are new. A = S(O)m, O or NR5; m = 0, 1 or 2; R5 = H, 1-8C alkyl or CmH2mR81; R81 = 3-8C cycloalkyl, phenyl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe, and NR82R83) or 1-9C heteroaryl (bonded via C or N and opt. substd. by 1-3 of F, Cl, CF3, Me, OMe, OH, NH2, NHMe and NMe2); R82, R83 = H or Me; one of R1, R2 = CO-N=C(NH2)2; the other = H, F, Cl, Br, I, 1-3C alkyl, OR6, 1-4C perfluoroalkyl, CO-N=C(NH2)2 or NR6R7; R6, R7 = H or 1-3C alkyl; R3, R4 = (i) H, F, Cl, Br, I, CN, X(CH2)m(1-6C) perfluoroalkyl, X(CH2)mF, S(O)mR8, CONR9R10, COR11, SO2NR12R13; (ii) 1-8C alkyl, CmH2mR81; (iii) 1-9C heteroaryl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe, OH, NH2, NHMe and NMe2); (iv) -Y-C6H4-(CO)i-(CHOH)j-(CHOH)k-R23; (v) H, F, Cl, Br, I, CN, 1-8C alkyl, 1-8C perfluoroalkyl, 3-8C alkenyl, CgH2g-R26; SR29, OR30, NR31R32, CR33R34R35; (vii) -W-C6H4-R97; (viii) S(O)mR37, SO2NR38R39; (ix) X1R46; (x) SR64, OR65, NHR66, NR67R68, CHRR69R70, CR54R55-OH, Ctriple bondC-R56 C(R58) C-R57 (sic), (CR59R60)u-CO-(CR61R62)v-R63; (xi) SO2NHR76; or (xii) NR84R85; X = O, S or NR14; R14 = H or 1-3C alkyl; R8 = 1-5C alkyl, 3-6C alkenyl, CnH2nR15 or CF3; R9, R11, R12 = H or as R8; n = 0-4; R15 = 3-7C cycloalkyl or phenyl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe and NR16R17); R16, R17 = H or 1-4C alkyl; R10, R13 = H or 1-4C alkyl; or R9+R10 or R12+R13 = (CH2)4 or (CH2)5 in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl; R18 = 3-8C cycloalkyl or phenyl (opt. substd. by 1-3 of F, Cl, CF3, Me, OMe and NR19R20); R19, R20 = H or Me; Y = O, S or NR22; h = 0 or 1; i, j, k = 0-4; provided that h, i and k are not all 0; R22, R23 = H or 1-3C alkyl; g = 0-4; R26 = 3-8C cycloalkyl, phenyl, biphenyl, or naphthyl (where aromatics are opt. substd. by 1-3 of F, Cl, CF3, Me, OMe and NR27R28); R27, R28 = H, 1-4C alkyl or 1-4C perfluoroalkyl; R29-R31, R33 = -(CH2)m- (1-9C) heteroaryl (opt. substd. as in R81); R32, R34, R35 = H, 1-4C alkyl, 1-4C perfluoroalkyl, or as R29; R96 = heteroaryl as defined for R81, or benzyl; W = O, S or NR36; R36 = H or 1-4C alkyl; R37 = 1-8C alkyl, 1-8C perfluoroalkyl, 3-8C alkenyl or -CsH2s-R40; s = 0-4; R40 = as R26; R38 = H, 1-8C alkyl, 1-8C perfluoroalkyl, 3-8C alkenyl or -CwH2w-R26; R39 = H, 1-4C alkyl or 1-4C perfluoroalkyl; or R38+R39 = (CH2)4 or (CH2)5, in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl; X1 = O, S, NR47, (D=O)A'- or NR48C=MN*(R49)-; M = O or S; A' = O or NR50; D = C or SO; R46, R49 = 1-8C alkyl, 3-8C alkenyl, -(CH2)b-(1-7C)perfluoroalkyl or -CxH2x-R26; b = 0 or 1; x = 0-4; R47, R48, R50 = H, 1-4C alkyl or 1-4C perfluoroalkyl;

or R46+R47 or R46+R48 = (CH2)4 or (CH2)5 in which CH2 may be replaced by O, S, NH, NMe or N-benzyl; A' and N* are bonded to the phenyl ring of the benzoylguanidine structure; R64-R67, R69 = -(CH2)y-(CHOH)z-(CH2)q'-(CH2OH)t-R71 or -(CH2)b'-O-(CH2CH2O)c'-R72; R71, R72 = H or Me; b', c' are not defined; u, t = 1-4; v, y, z, a' = 0-4; R68, R70, R54, R55 = H or 1-6C alkyl; or CR69R70 or CR54R55 = 3-8C cycloalkylidene; R63 = H, 1-6C alkyl, 3-8C cycloalkyl or -CeH2e-R73; e = 0-4; R80 = 5-7C cycloalkyl or phenyl (opt. substd. by 1-3 of F, Cl, CF3, OMe and 1-4C alkyl); or R77+R78 = (CH2)4 or (CH2)5, in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl; R79 = as R77; or amidino; R84, R85 = H or 1-4C alkyl; or R84+R85 = (CH2)4 or (CH2)5 in which one CH2 may be replaced by O, S, NH, NMe or N-benzyl or 1 or 2 CH2 gps. may be replaced by CH-Cd'H2d'+1; d' is not defined. Cpds. (I; A = O; R1 = -CON=C(NH2)2; R2, R3 = H; R4 = H, Me or Et) are excluded.

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Derwent Class: B03

International Patent Class (Main): C07D-000/00; C07D-207/34; C07D-207/40; C07D-207/416

International Patent Class (Additional): A01N-001/02; A61K-031/33; A61K-031/34; A61K-031/38; A61K-031/40; A61K-031/415; A61K-031/44; A61K-031/445; A61K-031/47; A61K-049/00; C07D-307/68; C07D-333/38; C07D-333/48; C07D-401/00; C07D-401/04; C07D-401/12; C07D-403/02; C07D-403/04; C07D-403/12; C07D-405/02; C07D-405/04; C07D-405/12; C07D-409/02; C07D-409/04; C07D-409/12; C07D-521/00

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New and known thienyl urea or isourea derivs. - used as animal growth promoters

Patent Assignee: BAYER AG (FARB)

Inventor: BERSCHAUER F; DEJONG A; HALLENBACH W; LINDEL H; SCHEER M

Number of Countries: 019 Number of Patents: 013

Patent Family:

| Patent No | Kind | Date | Applicat No | Kind | Date | Main IPC | Week |
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